

# One-Step Stain Free PAGE Gel Rapid Preparation Kit

QUICK START GUIDE

Research use only. Not for diagnostic procedures.

### **Contents** and storage

Product	Cat. No.	Quantity	Component	Cat. No.	Volume
	RF1521	100 Assays	Stacking Buffer A	RF1501	80 mL
One-Step Stain Free PAGE Gel Rapid Preparation Kit (6%)			Stacking Buffer B	RF1502	80 mL
			Resolving Buffer A (6%)	RF1503	250 mL
			Resolving Buffer B (6%)	RF1508	250 mL
			Enhanced Catalyst	RF1500	8 mL
One-Step Stain Free PAGE Gel Rapid Preparation Kit (7.5%)	RF1522	100 Assays	Stacking Buffer A	RF1501	80 mL
			Stacking Buffer B	RF1502	80 mL
			Resolving Buffer A (7.5%)	RF1504	250 mL
			Resolving Buffer B (7.5%)	RF1509	250 mL
			Enhanced Catalyst	RF1500	8 mL
One-Step Stain Free PAGE Gel Rapid Preparation Kit (10%)	RF1523	100 Assays	Stacking Buffer A	RF1501	80 mL
			Stacking Buffer B	RF1502	80 mL
			Resolving Buffer A (10%)	RF1505	250 mL
			Resolving Buffer B (10%)	RF1510	250 mL
			Enhanced Catalyst	RF1500	8 mL
	RF1524	100 Assays	Stacking Buffer A	RF1501	80 mL
			Stacking Buffer B	RF1502	80 mL
One-Step Stain Free PAGE Gel Rapid Preparation Kit (12.5%)			Resolving Buffer A (12.5%)	RF1506	250 mL
Rapid Preparation Kit (12.5%)			Resolving Buffer B (12.5%)	RF1511	250 mL
			Enhanced Catalyst	RF1500	8 mL
	RF1525	100 Assays	Stacking Buffer A	RF1501	80 mL
			Stacking Buffer B	RF1502	80 mL
One-Step Stain Free PAGE Gel Rapid Preparation Kit (15%)			Resolving Buffer A (15%)	RF1507	250 mL
			Resolving Buffer B (15%)	RF1512	250 mL
			Enhanced Catalyst	RF1500	8 mL

**Storage:** The enhanced catalyst should be stored at -20°C; other components are stored at 4°C for 12 months. Product shipped at ambient temperature. Once opened, the enhanced catalyst can be stored at 4°C for three months.

## Introduction

This kit simplifies polyacrylamide gel preparation for standard Tris-glycine electrophoresis. These solutions are formulated to allow for a quick casting method with **no polymerization wait time** between pouring the resolving and stacking gels. The stacking gel is color-coded (red, blue, or green) for easy sample loading.

Instead of TEMED, an odorless catalyst ensures rapid polymerization (15-30 minutes at room temperature). This versatile kit produces gels suitable for both denaturing and native PAGE, with **protein bands visualized** directly under UV light (302 nm).

## **Quick Cast Protocol**

#### Preparation of resolving solutions and stacking solutions for gel casting

For 1 mm thick mini-gel										
Gel Percentage	Resolving Buffer A	Resolving Buffer B	Catalyst	Stacking Buffer A	Stacking Buffer B	Catalyst				
Each	2.3 mL	2.3 mL	50µL	0.75 mL	0.75 mL	15µL				
For 0.75 mm thick mini-gel										
Gel Percentage	Resolving Buffer A	Resolving Buffer B	Catalyst	Stacking Buffer A	Stacking Buffer B	Catalyst				
	-	-	<b>Catalyst</b> 35µL	-	-	<b>Catalyst</b> 10µL				
Percentage	Buffer A	Buffer B		<b>Buffer A</b> 0.5 mL	Buffer B					

\*Volumes listed are sufficient for casting one 7.4 x 8.2 cm mini-gel and can be multiplied by N (desired number of gels) to cast multiple gels at once.

70µL

1.00 mL

1.00 mL

20µL

3.45 mL

3.45 mL

Each

#### Instructions:

- 1. Prepare the resolving gel with the desired acrylamide percentage by pipetting **equal volumes** of <u>Resolving Buffer A</u> and R<u>esolving Buffer B</u> into a clean conical tube.
- 2.Prep<mark>are the</mark> stacking gel by pipetting **equal volumes** of <u>Stacking Buffer A</u> and <u>Stacking Buffer B</u> into a clean conical tube.
- 3.Add the required volume of <u>Enhanced Catalyst</u> into the resolving tube. Gently mix reagents, avoiding the introduction of air bubbles into the gel mixture.Using a pipette, fill each cassette to 0.5-1 cm below the comb teeth.
- 4.Add the required volume of <u>Enhanced Catalyst</u> into the stacking tube. Gently mix reagents, avoiding the introduction of air bubbles into the gel mixture.
- 5. Position the pipette at the middle of the cassette and gently add the stacking gel, filling to the top of the short plate. A dip may occur where pipetting takes place but will level out.
- 6.Quickly and carefully insert the comb and avoid air bubble entrapment below the teeth.
- 7.Allow gels to polymerize for 15 minutes.
- 8.Gels can be used immediately or wrapped in DI water-soaked paper towels and stored in an airtight container at 4°C for up to 5 days.
- 9. Remove the comb before loading samples. The recommended electrophoresis conditions are **150V for 60 minutes** or **200V for 45 minutes**.
- 10. Following electrophoresis, carefully remove the gel cassette and glass slides. Protein bands can be visualized under **UV light** (302 nm).
- 11. After transferring proteins to an NC or PVDF membrane, expose it again to UV light to confirm protein transfer.
- 12. Please note that the excitation of fluorescent dyes may take approximately 1-5 minutes. Please be patient until the protein bands become clearly visible.
- 13. For optimal visualization of fluorescent protein bands on the membrane, preexposure to UV light may be necessary before protein transfer. This preexcitation step can enhance fluorescence signal intensity.

Note:

1. Gel polymerization time may vary depending on temperature. Higher temperatures generally accelerate polymerization, while lower temperatures may prolong the process. In colder environments, consider increasing the Catalyst volume by up to twofold.

2. If the kit has been stored at 4°C, allow it to reach room temperature before use to minimize the formation of air bubbles during gel casting.

